A controlled study of dementia in Parkinson's disease

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SUMMARY Tests of cognitive functions were carried out in a group of patients with Parkinson's disease and repeated after a three-year interval. Comparison was made with a control group drawn from a population of psychiatric patients, matched for age and sex. No differences in cognitive functions were found between the groups, either initially, or between those surviving for three years. Deaths among the index group included a high proportion of patients with cognitive impairment and there was an increasing prevalence and severity of dementia in the index group which exceeded that observed in the control group. Requirements for a methodologically sound study of dementia in Parkinson's disease are discussed.

Since James Parkinson's original description of paralysis agitans many have questioned his statement "the senses and intellect are uninjured" (1817).1 Ball was among the first to suggest that intellectual impairment may occur in Parkinson's disease: "paralysis agitans is accompanied more often than is thought by intellectual difficulties" (1882).2 Dementia in Parkinson's disease has been estimated to occur in between 14% and 40% of those affected by the disorder.3-7 Dementia has been described as occurring particularly in the "so-called" arteriosclerotic type of Parkinsonism8 but a retrospective study conducted more recently showed evidence of dementia in one third of patients with only minor differences in the proportions of the three main aetiological types: idiopathic Parkinson's disease, postencephalitic Parkinsonism and arteriosclerotic Parkinsonism.9

In studies which have used control groups for comparison and standardised methods of assessment of intellectual function the evidence for dementia being a regular feature of Parkinson's disease is less impressive. At least two well-conducted studies have failed to show evidence of dementia in Parkinson's disease.10,11 Others have agreed with these workers, suggesting that any apparent decline in mental function could be attributed to psychomotor slowing,12 or alternatively to the effects of depression of mood.7 Against this view, some authors have concluded that there is a "widely generalised deterioration" apart from that in motor function or that there is an associated memory impairment in Parkinson's disease.13,14 In a controlled follow-up study of a large number of subjects 32% were found to show evidence of dementia and this was ten times the prevalence of dementia among their spouses.15 Other evidence is interpreted as suggesting the presence of specific deficits in Parkinson's disease, that these are minor as compared with the motor changes seen, and often static, and that to apply the label of dementia is both "unacceptable and misleading".16 Other workers have suggested that the deficit is similar to that seen in patients following frontal lobectomy, although part of their psychological assessment did suggest impairment of short-term memory and attention.17,18 An important suggestion is that patients with Parkinson's disease, among others, suffer from a distinct type of dementia attributable to disease of the basal ganglia rather than of the cortex.19 This syndrome, known as "subcortical dementia", is characterised by: amnesia, impaired ability to manipulate acquired knowledge, slowness of the thought processes, and changes in personality. Although a clinical picture resembling this description may be seen in Parkinson's disease, the status of "sub-cortical dementia" remains unclear.19,20,21

The uncertainty which remains over this important issue led us to carry out a further investigation. Our study is based on two hypotheses: firstly, if dementia is a clinical feature of Parkinson's disease rather than a chance finding, it will be more common among patients with the disease than in a control group; and secondly, if dementia is an integral part of the syndrome, progression of the disease will lead to an
increasing prevalence and severity of dementia among sufferers.

**Method**

Forty patients suffering from Parkinson's disease (idiopathic in thirty-seven and postencephalitic in three) were compared with a control group matched for age and sex. The patients suffering from Parkinson's disease were attending a neurological out-patient clinic and the diagnosis was confirmed by two neurologists. The patients continued to receive anti-Parkinson medication.

**Table 1 Male index patients with their matched controls**

<table>
<thead>
<tr>
<th>Parkinson patients</th>
<th>Control group</th>
<th>Initial psychiatric diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yr)</td>
<td>Intellectual rating</td>
<td>Year 0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>58</td>
<td>1</td>
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<td>55</td>
<td>0</td>
<td>Died</td>
</tr>
<tr>
<td>67</td>
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<td>Not Ass'd</td>
</tr>
<tr>
<td>64</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>63</td>
<td>2</td>
<td>2</td>
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<td>59</td>
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<tr>
<td>74</td>
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<tr>
<td>69</td>
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<td>Died*</td>
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<tr>
<td>77</td>
<td>0</td>
<td>Died</td>
</tr>
<tr>
<td>77</td>
<td>3</td>
<td>Died*</td>
</tr>
<tr>
<td>61</td>
<td>0</td>
<td>Died</td>
</tr>
<tr>
<td>65</td>
<td>0</td>
<td>Not Ass'd ‡</td>
</tr>
</tbody>
</table>

* Patients initially demented who died.
† Patients who became demented during follow-up.
‡ Patients who showed progression of dementia.
? Not fully assessed.

**Table 2 Female index patients with their matched controls**

<table>
<thead>
<tr>
<th>Parkinson patients</th>
<th>Control group</th>
<th>Initial psychiatric diagnosis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (yr)</td>
<td>Intellectual rating</td>
<td>Year 0</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>62</td>
<td>1</td>
<td>Not Ass'd</td>
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<td>74</td>
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<td>0</td>
</tr>
<tr>
<td>68</td>
<td>0</td>
<td>1†</td>
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<tr>
<td>61</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>53</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>70</td>
<td>0</td>
<td>0</td>
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<td>64</td>
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<td>1†</td>
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<td>63</td>
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<td>2†</td>
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<tr>
<td>64</td>
<td>1</td>
<td>2†</td>
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<td>63</td>
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<tr>
<td>65</td>
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<td>2†</td>
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<td>68</td>
<td>1</td>
<td>2†</td>
</tr>
<tr>
<td>62</td>
<td>3</td>
<td>Died*</td>
</tr>
</tbody>
</table>

* Patients initially demented who died.
† Patients who became demented during follow-up.
‡ Patients who showed progression of dementia.

The control group was selected retrospectively from the outpatient practices of the four consultant psychiatrists who together formed the senior medical staff of the clinical unit of the Department of Psychiatry, Nottingham University Medical School. A control group of this kind was chosen for two reasons: first, we chose to use a "patient" rather than a "normal" control group because there is evidence that any kind of illness may impair performance in tests which measure cognitive function, (D Bannister, personal communication), and we wished to compare two ill groups; and second, it was essential to have control subjects on ...
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whom full details of cognitive function had been recorded.

The staff of the records department was asked to give the
names of patients who had been seen in the out-patient
department during a period two to three years previously.
Each index patient was individually matched with a control
patient by sex and age; this meant that if an index patient
was lost to the study, the corresponding control patient was
also excluded from comparisons with the survivors of the
index group. Because of the relatively narrow age range of
the index group, the gathering of the control group was a
laborious task. The outcome of the matching procedure
is shown in tables 1 and 2. Once the control group was
identified, arrangements were made to reassess the patients
on or near the third anniversary of their first out-patient
attendance. One of the subjects chosen as a control refused
reassessment which left one index patient without a control
pairing. The average age of the control group is higher than
of the index group: this would increase any psychiatric
morbidity closely linked with age in the control group and
would be likely to reduce the contrasts between the groups
in certain respects.

The index group were assessed using a small battery of
tests of cognitive function as part of a more comprehensive
study.22 They were reassessed using the same battery three
years later. The control group were assessed using the same
standardised tests at follow-up and their scores three years
previously were calculated by scrutiny of the notes recorded
by the consultant psychiatrist at the time of their first
assessment.

The tests of cognitive function are derived from those
used in ordinary clinical practice and include tests of
orientation, sentence-learning, counting backwards,
counting by threes, digit retention, five-minute memory, a
paired word-learning test, simple arithmetic and general
knowledge. These tests, which do not include tests of motor
skills, were performed in a standardised way and led to
each patient receiving a score of intellectual impairment
according to the General Practice Research Unit Interview
Schedule (1970).23

Results

The index group of forty patients, comprising twenty-
three men and seventeen women had an average age
of 61·9 years. (Average length of illness 9 years,
range 1–42 years. 30 on levodopa, six had had neuro-
surgical treatment.) The average age of the control
group was 65·7 years.

Initial Assessment  The results of the initial assess-
ment of intellectual function are shown in table 3. If
all the patients showing some evidence of intellectual
impairment are grouped together irrespective of severity, 40% of the index group and 32% of the
control group are seen to be affected. This difference
does not reach statistical significance ($X^2 = 0.747$,
$p = 0.20$, df = 1). When only those patients showing
intellectual impairment of a degree which is definitely
pathological (that is, groups 2, 3 and 4) are compared,
20% of the index group and 24% of the control group

<table>
<thead>
<tr>
<th>Initial assessment of intellectual function</th>
<th>Index group</th>
<th>Control group</th>
</tr>
</thead>
<tbody>
<tr>
<td>No abnormality</td>
<td>0</td>
<td>27</td>
</tr>
<tr>
<td>Mild impairment</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td>Definitely pathological</td>
<td>2</td>
<td>8</td>
</tr>
<tr>
<td>Moderate impairment</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>Marked impairment</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Totals</td>
<td>40</td>
<td>39</td>
</tr>
</tbody>
</table>

(show these levels of impairment; again the difference
between the groups is not statistically significant ($X^2$
$= 0.18$, $p = 0.5$, df = 1).

Assessment after three years The results of the
assessment: after three-year interval were again
compared in the index and control groups. Ten
patients were lost from the index group. The
survivors were compared with their corresponding
controls (table 4). Thirty-three per cent of the index
group showed some impairment as compared with
27% of the control group; this difference is not sta-
tistically significant ($X^2 = 0.314$, $p = 0.5$, df = 1). When
those subjects with “definitely pathological” impair-
ment are compared, there are 26% in the index group
compared with 20% in the control group; this difference
is not statistically significant ($X^2 = 1.14$, $p = 0.2$, df = 1).

Patients lost at three years Of the ten patients lost to
the study seven were known to have died, two were
not formally assessed and one was untraceable. The two
patients not formally assessed were still in contact
with the clinic; one was so disabled that she was
unable to answer the door and the interviewer was
unaware that he should put his hand through the
letter box and pull out the key which was on a string;
the other patient, who was well-known to the
interviewer, came to the door, did not recognise him,
and spoke in a way which showed him to be
disorientated and disinhibited, saying “We must meet
for a drink another time, it's not convenient now”.
Thus, there is definite knowledge of death in seven
subjects of whom four are known to have been

<table>
<thead>
<tr>
<th>Assessment of intellectual function after three years</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intellectual rating</td>
</tr>
<tr>
<td>--------------------</td>
</tr>
<tr>
<td>0</td>
</tr>
<tr>
<td>1</td>
</tr>
<tr>
<td>2</td>
</tr>
<tr>
<td>3</td>
</tr>
<tr>
<td>4</td>
</tr>
<tr>
<td>Totals</td>
</tr>
</tbody>
</table>

(Percentages in brackets)
demented, evidence of dementia in one still living and severe disability in another.

Morbidity from dementia  In the index group the proportion of subjects initially rated as definitely impaired and surviving to be examined after 3 years was only 13%. The ten patients lost from the index group were worse than average for the group in terms of intellectual impairment; of the initial assessment 50% were judged to have some degree of intellectual impairment; 10% mild, 20% moderate and 20% marked. Four of those who died had dementia, and a patient lost to the study but not known to have died was demented at the initial assessment.

Four (including one not fully assessed) index patients showed evidence of dementia for the first time during the follow-up period, as compared with one subject in the control group. Three patients in the index group showed evidence of dementia initially, but this disappeared by the time of the follow-up assessment three years later: this change is likely to be an effect of treatment with levodopa.

Five patients in the index group showed progression in the severity of dementia during the three year period as compared with two patients in the control group.

The details of patients studied and of their controls is shown in tables 1 and 2 and the various indices of morbidity are summarised in table 5. The numbers of patients involved were so small and the changes observed so diverse as to prevent statistical evaluation of most of the follow-up data.

Discussion

Although the index and control groups were similar as regards the prevalence of dementia initially, by the end of the three year follow-up period, evidence had accumulated which demonstrates a higher morbidity from dementia among the patients with Parkinson’s disease. This was shown in three ways: patients suffering from dementia were more likely to die during the period of study, only 13% surviving 3 years: dementia observed for the first time during the follow-up period was more common in the index group; there was more evidence of progression in the severity of dementia in the index group. On these findings we are unable to reject either of our initial hypotheses. Within the methodological limits of this small scale study it seems likely that dementia occurs more commonly in Parkinson’s disease than in a psychiatric population, and should be regarded as an intrinsic part of the disorder as it occurs with increasing prevalence and severity as the disease progresses. These conclusions must be qualified and do not constitute final evidence of the status of dementia in Parkinson’s disease: a larger scale study with certain improvements in method is required for the matter to be finally clarified.

The incidence of dementia in our index group initially was 20%; this is a low figure by comparison with some other studies. Celesia and Wanamaker believed that 40% of their patients with Parkinson’s disease showed some evidence of dementia. Martilla and his colleagues found an overall incidence of dementia of 28.8%. Pollack and Hornabrook, found significant mental deterioration in 20% and Hoehn and Yahr found an incidence of “mild to moderate organic mental syndrome” in 14%. These studies did not, however, involve comparison with control populations. Loranger and his colleagues, using the Wechsler Adult Intelligence Scale on a patient group alone, found impairment in 36.5% and conclude that the intellectual decline is twice that of the normal population of the same age. Lieberman and his colleagues found 32% of 520 patients to have impaired cognitive functioning and that this was ten times the prevalence of dementia among 407 spouses. These workers and others have suggested that there is a sub-group of patients with Parkinson’s disease in which dementia is a prominent feature. Our results are compatible with this suggestion without providing positive evidence in support. There is some pathological data which suggests a relationship between the dementia of Parkinson’s disease and that of Alzheimer’s disease.

The low incidence of dementia in our patients as compared with other studies and the loss of demented patients due largely to death, questions the adequacy of the method of our study in bringing out any difference which might exist between the index and control groups. Two aspects of the method of the study require special scrutiny: first, was the ‘type of control group appropriately chosen; and secondly, were adequate methods used in assessing intellectual function?

The control group was taken from a large population of patients who had seen a psychiatrist at least once. This group of patients might be expected to contain some patients suffering from dementia and
probably a higher proportion than would be found in the general population. In this way the composition of the control group would reduce any contrast between the index group and the control group with respect to dementia. On the other hand, the presence of an illness of any kind is known to impair performance on tests of intellectual function in a non-specific way so that the choice of a “patient” control group was more appropriate than a group of normal subjects. Our experience suggests that a control group drawn from non-psychiatric patients would have been more satisfactory, probably leading to a reduction in the proportion of patients suffering from dementia and thereby sharpening any contrast with the index group. A “spouse” control group has been used in some studies and has proved to be fairly satisfactory.15 There are, however, several problems with this approach: not all patients have spouses; spouses tend to be younger than patients; spouses are generally a “well” group and would be expected to show less cognitive impairment than a “patient” group. (D Bannister, personal communication).

Our control group was less satisfactory than it might have been because it was selected retrospectively; this meant that the initial assessment was carried out by a clinician for purely clinical purposes without using the standardised assessment subsequently used in follow-up. This method of choosing the control group prevented comparison of death rates in the groups as selection meant that the control subjects had already survived the follow-up period. A control group studied prospectively and in parallel with the index group would be more satisfactory but would require a large scale study.

There are particular problems in assessing cognitive function in patients suffering from Parkinson’s disease and these have to be considered in making a choice of methods of assessment.29 Patients are usually slow both in action and in thought and often have difficulty in sustaining concentration. Furthermore, it is important to use tasks which all subjects in a study are capable of completing on each occasion on which they are administered. A compromise has to be made between rigour and practicability. For these reasons we chose a relatively short and simple battery of tests which included none which depend upon motor skills. This battery is not refined and has not been subjected to rigorous tests of reliability; it was, however, administered in a standardised way and was found to be acceptable to patients and understood by them. In spite of the choice of a relatively simple battery of tests some patients required 45 minutes to complete it; this must be close to the limit of concentrated application in such a group of patients. The methods of assessment used in the well controlled and informative study by Lieberman and his colleagues, were of a similar kind and have the same strengths and weaknesses.15

A matter which requires comment is the possible effect of treatment with levodopa on performance in tests of cognitive functions. A number of studies have reported improvement in intelligence, perceptual, motor and memory functions, following administration of levodopa. These effects are more limited when treatment and observations are continued over a period of years.28 Various explanations of the effect of levodopa on cognitive functions have been put forward; but the most favoured is that the effect is mediated through enhanced arousal or activation in a non-specific way.30

Our study, which is methodologically different from studies previously performed, brings further evidence which suggests that dementia is a regular feature of Parkinson’s disease. This evidence, and the evidence from previous studies, is not, however, conclusive. Our study highlights some of the methodological problems in investigating dementia in Parkinson’s disease, and it would appear that a completely satisfactory study has not yet been completed. Our experience underlines the importance of an appropriately selected control group, the use of standardised and appropriate methods of assessment which can be successfully carried out in patients with Parkinson’s disease, the study of patients over a suitable period of time with measures which allow for the loss of patients from relevant causes, and numbers of subjects and controls which would allow full statistical evaluation of the results. The possible existence of a sub-group of patients with Parkinson’s disease in which dementia is a regular feature, and the status of the concept of “sub-cortical dementia” deserve further investigation.

We are indebted to the following medical colleagues: Professor CD Marsden, Dr JD Parkes and Dr KJ Zilkha of King’s College Hospital for allowing us to study patients suffering from Parkinson’s disease who were attending their out-patient clinics, Professor JE Cooper, Dr IB Pearson and Dr DA Toms of Mapperley Hospital, Nottingham, for allowing us to select the control group from their patients.

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